(19) World Intellectual Property Organization International Bureau





(43) International Publication Date 17 April 2003 (17.04.2003)

PCT

(10) International Publication Number WO 03/031632 A1

(51) International Patent Classification7:

C12N 15/82

- (21) International Application Number: PCT/EP02/11188
- (22) International Filing Date: 2 October 2002 (02.10.2002)
- (25) Filing Language:

English

(26) Publication Language:

English

- (30) Priority Data: 01203760.2 5 October 2001 (05.10.2001) E
- (71) Applicant (for all designated States except US): VLAAMS INTERUNIVERSITAIR INSTITUUT VOOR BIOTECHNOLOGIE VZW [BE/BE]; Rijvisschestraat 120, B-9052 Zwijnaarde (BE).
- (72) Inventors; and
- (75) Inventors/Applicants (for US only): DEPICKER, Anna [BE/BE]; Molenstraat 61, B-9820 Merelbeke (BE). VAN HOUDT, Helena [BE/BE]; Lieskensweg 7, B-9080 Lochristi (BE).
- (74) Common Representative: VLAAMS INTERUNIVER-SITAIR INSTITUUT VOOR BIOTECHNOLOGIE VZW; Rijvisschestraat 120, B-9052 Zwijnaarde (BE).

- (81) Designated States (national): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW.
- (84) Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, SK, TR), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

Published:

with international search report

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

3/031632 A1

(54) Title: AN EFFICIENT SYSTEM FOR RNA SILENCING

(57) Abstract: The invention relates to a method for efficient RNA silencing of target genes in eucaryotic cells, particularly plant cells. Consequently, the method can be used to reduce the phenotypic expression of an endogenous gene in a plant cell. Furthermore the method can be applied in a high throughput screening for mutant phenotypes as a result of RNA silencing of any endogene.

An efficient system for RNA silencing

10

15

20

25

30

Field of the invention

The invention relates to a method for efficient RNA silencing in eucaryotic cells, particularly plant cells. Consequently, the method can be used to reduce the phenotypic expression of an endogenous gene in a plant cell. Furthermore the method can be applied in a high throughput screening for RNA silencing.

Background of the invention

RNA silencing is a type of gene regulation based on sequence-specific targeting and degradation of RNA. The term encompasses related pathways found in a broad range of eukaryotic organisms, including fungi, plants, and animals. In plants, RNA silencing serves as an antiviral defense, and many plant viruses encode suppressors of silencing. Also it becomes clear that elements of the RNA silencing system are essential for gene regulation in development. The emerging view is that RNA silencing is part of a sophisticated network of interconnected pathways for cellular defense, transposon surveillance, and regulation of development. Based on the sequence specific RNA degradation, RNA silencing has become a powerful tool to manipulate gene expression experimentally. RNA silencing was first discovered in transgenic plants, where it was termed co-suppression or posttranscriptional gene silencing (PTGS). Sequence-specific RNA degradation processes related to PTGS have also been found in ciliates, fungi, and a variety of animals from Caenorhabditis elegans to mice (RNA interference). A key feature uniting the RNA silencing pathways in different organisms is the importance of double-stranded RNA (dsRNA) as a trigger or an intermediate. The dsRNA is cleaved into small interfering RNAs (21 to 25 nucleotides) of both polarities, and these are thought to act as guides to direct the RNA degradation machinery to the target RNAs. An intriguing aspect of RNA silencing in plants is that it can be triggered locally and then spread via a mobile silencing signal. In plants, RNA silencing is correlated with methylation of homologous transgene DNA in the nucleus. Other types of epigenetic modifications may be associated with silencing in other organisms.

It is known from the art that transgenes encoding ds or self-complementary (hairpin) RNAs of endogenous gene sequences are highly effective at directing the cell's degradation mechanism against endogenous (ss) mRNAs, thus giving targeted gene

5

10

15

20

25

30

suppression. This discovery has enabled the transgenic enhancement of a plant's defense mechanism against viruses that it is unable to combat unaided. It has also shed light on how antisense and co-suppression might operate: by the inadvertent integration of two copies of the transgenes in an inverted repeat orientation, such that read-through transcription from one gene into the adjacent copy produces RNA with self-complementary sequences.

RNA silencing is induced in plants by transgenes designed to produce either sense or antisense transcripts. Furthermore, transgenes engineered to produce self-complementary transcripts (dsRNAs) are potent and consistent inducers of RNA silencing. Finally, replication of plant viruses, many of which produce dsRNA replication intermediates, causes a type of RNA silencing called Virus Induced Gene Silencing (VIGS). Whether VIGS, and the different types of transgene-induced RNA silencing in plants result from similar or distinct mechanisms is still a matter of debate. However, recent genetic evidence raises the possibility that the RNA silencing pathway is branched and that the branches converge in the production of dsRNA.

Until recently RNA silencing was viewed primarily as a thorn in the side of plant molecular geneticists, limiting expression of transgenes and interfering with a number of applications that require consistent, high-level transgene expression. With our present understanding of the process, however, it is clear that RNA silencing could have enormous potential for engineering control of gene expression, as well as for the use as a tool in functional genomics. It could be experimentally induced and targeted to a single specific gene or even to a family of related genes. Likewise, ds RNAinduced TGS may have similar potential to control gene expression. Although several methods for RNA silencing have been described in the art (WO99/53050, WO99/32619, WO99/61632, and W098/53083), there is clearly a need to develop alternative and more efficient tools for RNA silencing. In the present invention we have developed a highly efficient method for RNA silencing that can also be used as a tool for high throughput silencing. Said method uses a host that carries already a silenced locus and a second recombinant gene comprising a region that is homologous with the silenced locus. Although it is known from the art that the recombinant gene will be silenced, we have surprisingly found that also target genes, which have no significant homology with the silenced locus but have homology with the recombinant gene, are efficiently silenced.

Figure legends

Fig. 1: Schematical outline of homology between a silenced locus X, a recombinant gene Y and a target gene Z.

Fig. 2: Schematical outline of the T-DNA constructs that are present in silenced locus X₁, recombinant gene Y₁ and target gene Z₁ (T-DNAs of pGVCHS287, pGUSchsS and pXD610 respectively) and of the transcript homology between X₁, Y₁ and Z₁.

LB and RB: left and right T-DNA border respectively; Pnos: nopaline synthase promoter; hpt: hygromycin phosphotransferase coding sequence; 3'nos: 3'untranslated region of the nopaline synthase gene; P35S; Cauliflower mosaic virus 35S promoter; nptll c.s., neomycin phosphotransferase II coding sequence; 3'chs: 3'untranslated region of the chalcone synthase gene of Anthirrinum majus; +1: transcription start; An: poly A-tail; gus c.s.: β-glucuronidase coding sequence; Pss: promoter of the small subunit of rubisco; bar: phosphinotricine transferase coding sequence; 3'g7: 3'untranslated region of the Agrobacterium octopine T-DNA gene 7; 3'ocs: 3'untranslated region of octopine synthase gene.

Fig. 3: Schematical outline of the T-DNA construct present in silenced locus X₁ and of the transiently introduced T-DNAs Y₂ (T-DNAs of pGVCHS287 and pPs35SCAT1S3chs, respectively) and of the transcript homology between X₁, Y₂ and Z₂ (the catalase1 endogene). Abbreviations as in Fig. 2

Fig. 4: Schematical outline of the T-DNA constructs present in silenced locus X₂ and of the transiently introduced T-DNAs Y₂ (T-DNAs of pGUSchsS + pGUSchsAS, and pPs35SCAT1S3chs, respectively) and of the transcript homology between X₂, Y₂ and Z₂ (the catalase1 endogene). Abbreviations as in Fig. 2

Fig.5: pPs35SCAT1S3chs

30

10

15

Detailed description of the invention

The present invention deals with an efficient method for RNA silencing in an eucaryotic host. The method makes use of a host that already comprises a silenced locus. Such a silenced locus can for example be generated by methods known in the art. For

example the publication of De Buck and Depicker, 2001 and other publications, and also patents WO99/53050, WO99/32619, WO99/61632, and W098/53083 describe methods to obtain RNA silencing and for generating a silenced recombinant locus. The 'target gene' is here defined as the gene of interest for silencing or to down-regulate its expression. An important aspect of this invention is that said target gene has no significant homology with the silenced locus. No significant homology means that either the overall homology is less than 40, 35, 30, 25% or even less, or that no contiguous stretch of at least 23 identical nucleotides are present (Thomas et al., 2001). Homology is typically measured using sequence analysis software (e.g., Sequence Analysis Software Package of the Genetics Computer Group, University of Wisconsin Biotechnology Center, 1710 University Avenue, Madison, Wis. 53705). Such software matches similar sequences by assigning degrees of homology to various insertions, deletions, substitutions, and other modifications. Silencing of said target gene in the present invention occurs via an intermediate step and hence our method is designated as domino silencing (Fig. 1). In said intermediate step a recombinant gene construct is introduced by transformation into the host comprising the silenced locus. Said recombinant gene construct has a region of homology with the silenced locus already present. Said region of homology is preferably more than 60, 70, 80, 90, 95 or even more than 99% homologous. The homologous region between the silenced locus and said recombinant gene can be found in the 5' untranslated or 3' untranslated region of the recombinant gene construct. Furthermore, said recombinant gene construct has a region of minimal 23 nucleotides (Thomas et al., 2001), but preferably longer, that are identical with the target gene, or has a region of overall homology of more than 60, 70, 80, 90, 95 or even more than 99%. A recombinant gene is defined herein as a construct which does not naturally occur in nature. A non-limiting example of a recombinant gene construct is a construct wherein the coding region of a gene is operably linked to a 5' untranslated region and/or to a 3' untranslated region of one or more other genes, alternatively said 5' or 3' untranslated region is an artificial sequence.

10

15

20

25

Thus in one embodiment the invention provides a method for obtaining efficient RNA silencing of a target gene comprising the introduction of a recombinant gene into a host that comprises a silenced locus and an unsilenced target gene whereby said recombinant gene comprises a region that is homologous with said silenced locus and

whereby said target gene has homology with said recombinant gene but has no significant homology with said silenced locus.

In another embodiment the method is used wherein said host is a plant or plant cell.

In another embodiment the method of the invention can be used for high throughput gene silencing. Indeed, a recombinant gene library can be made wherein for example every gene or coding region thereof is combined with (operably linked with) a region of homology with the silenced gene that resides in the silenced locus and said recombinant gene library can be transformed to an eukaryotic host or individual (specific) genes derived from said recombinant gene library can be transformed into an eukaryotic host wherein silencing of specific genes is wanted.

In yet another embodiment the invention provides a plant or plant cell that comprises a silenced locus and wherein a silenced target gene is obtained through the introduction of a recombinant gene according to the current method of the invention.

In yet another embodiment the RNA silencing of the target gene is obtained in more than 80, 85, 90 or 95% of the transgenic organisms.

In yet another embodiment the RNA silencing of the target gene occurs at an efficiency of more than 80, 85, 90 or 95 % as compared to the level of the unsilenced expression of the target gene.

20 Examples

5

10

15

25

30

A posttranscriptionally silenced inverted repeat transgene locus can trigger silencing of a reporter gene producing non-homologous transcripts.

We studied the interaction between three transgene loci X_1 , Y_1 and Z_1 (Fig. 2, For a detailed description of all loci and constructs, see materials and methods) to address the question whether or not a stepwise homology between loci can lead to silencing.

It has been demonstrated previously that the posttranscriptionally silenced nptII genes in locus X_1 are capable to in trans silence transiently expressed genes with partial transcript homology to their nptII transcripts (Van Houdt et al., 2000 b). We subsequently found that also a stably expressed β -glucuronidase (gus) gene (in locus Y_1), with partial transcript homology to the nptII transcripts of the silencing inducing locus X_1 , becomes efficiently silenced in trans (Fig. 2: X_1 and Y_1 and table 1: X_1Y_1 compared to Y_1). On the contrary, the nptII genes of locus X_1 are not able to trigger silencing of the gus genes in locus Z_1 which is expected as the genes of both loci produce transcripts without significant homology (Fig. 2). The homology between the

5

10

15

20

25

30

two transcripts of X_1 and Y_1 is mainly situated in the 3'untranslated region (250 nucleotides), but also the 5'untranslated sequences show a small region of homology (29 nucleotides). These results demonstrate that the in trans silencing effects are not triggered by promoter homology. When Y_1 and Z_1 loci are combined in so called Y_1Z_1 hybrids both types of gus genes, having transcript homology in the gus coding sequence of 1809 nucleotides, remain highly expressed as reflected in the normal gus activity showing that the RNA silencing mechanism does not become activated (Table 1: Y_1Z_1 compared to Y_1 and Z_1). Surprisingly, upon creation of a stepwise homology between X_1 and Z_1 by introducing locus Y_1 , the new observation described here is that also the gus expression in locus Z_1 is reduced in $X_1Y_1Z_1$ plants (Table 1: $X_1Y_1Z_1$ compared to Y_1Z_1). Thus, creating a stepwise homology between a silenced locus and a target gene by introducing a recombinant gene is sufficient to trigger silencing of the target.

Silencing inducing transgene loci can trigger silencing of a non-homologous endogene. We further assessed the universality and the usefulness in high throughput functional gene analyses of silencing elicited by a stepwise homology in trans, called domino silencing. Therefore, we evaluated whether the expression of the tobacco endogenous catalase1 (cat1) genes is reduced in plants carrying a silencing locus (X locus) showing no significant homology with the catalase endogene by introducing a recombinant gene (Y construct). As silencing locus we used either X₁ or X₂ (Fig.2: locus X₁, Fig.3: locus X₂), in either case containing the 3' chalcone synthase sequences of Anthirrinum majus (3'chs). As transmitter for silencing we constructed a recombinant gene composed of the catalase1 coding sequence and the 3' chs region under control of the 35S promoter (P35S) (residing on T-DNA pPs35SCAT1S3chs. Fig.2 and 3: T-DNA in Y₂). The recombinant cat1 3'chs genes (Y2) were introduced in tobacco leaves bearing locus X1 (or X2) via Agrobacterium injection. As a negative control, we introduced a recombinant gene in which the cat1 coding sequence is replaced by the gus coding sequence (pGUSchsS, T-DNA construct as in locus Y1 Fig.1). In this case, no stepwise homology is created between the silencing inducing locus and the target catalase endogenes. As a positive control, the recombinant construct Y2 was also introduced in transgenic tobacco with silenced catalase1 genes by the presence of a catalase1 antisense construct (Cat1AS in Champnongpol et al., 1996). Sixteen days after Agrobacterium injection, the catalase activity was determined

in protein extracts of injected leaf tissue and compared with the activity in non-injected wild type (SR1) leaf tissue (Table 2). The results indicate that domino silencing is also applicable to endogenes since the catalase activity is clearly reduced in 6 out of 7 samples, while it remains high in the negative controls. In conclusion, not only an inverted repeat-bearing silencing-inducing transgene locus, but also a silencing-inducing locus in which the two residing chimeric genes give rise to transcripts with complementarity in the 3'UTR (3'chs)(Fig.3: X₂), is able to trigger domino silencing reducing endogenous catalase expression.

10 <u>Table 1:</u> Results of a GUS-activity determination in protein extracts of leaf tissue harvested from tobacco plants containing different combinations of the loci X₁, Y₁ and Z₁ (Fig.2). The mean values of a number of plants (n) are given.

genotype	GUS-act. ¹ 4 weeks ²	N	GUS-act. Mature ²	n
	U GUS/mg TSP	}	U GUS/mg TSP	
X ₁	<3	1	<	1
Y ₁	368 ± 1654	9	n.d.	-
Z ₁	126 ± 30	10	48 ± 8	5
X ₁ Y ₁	2 ± 1	4	4 ± 2	4
X ₁ Z ₁	139 ± 35	9	46 ± 14	5
Y ₁ Z ₁	477 ± 101	10	231 ± 106	6
$X_1Y_1Z_1^5 \rightarrow Y_1Z_1$	195 ± 104	16	315 ± 46	8
$\rightarrow X_1Y_1Z_1$	4 ± 3	22	12 ± 4	9

¹ The mean GUS-activity (GUS-act.) was calculated, using n samples and expressed as units (U) GUS per milligram of total soluble protein (TSP).

² The plants were analyzed in two different developmental stages; 4 weeks after sowing and at a mature stage just before onset of flowering.

³ below detection limit (1 U GUS/mg TSP)

^{20 &}lt;sup>4</sup> standard deviation

⁵ Growth of $X_1Y_1Z_1$ plants was performed in conditions that both Y_1Z_1 and $X_1Y_1Z_1$ plants were able to develop. A PCR screen with X_1 -specific primers was performed to discriminate between presence and absence of X_1 .

n.d. not determined

<u>Table 2:</u> Results of a catalase-activity determination in protein extracts of leaf tissue harvested from Agrobacterium injected tobacco leaves.

Genotype injected	Construct introduced via	catalase activity 16 days
plant	Agrobacterium injection	after injection (60 μg TSP)
WT (SR1)	- (non-injected)	-0.2116 ² 100% ³
X ₁	PGUSchsS	-0.2556 121%
X ₁	Y ₂	-0.0589 27%
X ₁ ⁴	Y ₂	-0.0698 33%
X ₂	PGUSchsS	-0.1782 84%
X ₂	Y ₂	-0.0641 30%
X ₂	Y ₂	-0.0987 47%
X ₂ ⁴	Y ₂	-0.0914 43%
X ₂ ⁴	Y ₂	-0.1996 94%
X ₂ ⁴	Y ₂	-0.0627 30%
Cat1AS	Y ₂	-0.0439 21%

 $^{^{1}}$ X₁, see Fig. 3; X₂, see Fig. 4

⁴ 24 hours after Agrobacterium injection, the plants were placed under high light conditions for 24 hours (1000 μmol / m² s). This treatment is known to stimulate endogenous catalase 1 transcription. As the degree of cat suppression is similar in uninduced as in induced situation, the data indicate that enhanced transcription of the endogenous catalase target is not required to trigger domino silencing.

15

10

Materials and Methods

Plasmid construction

pPs35SCAT1S3chs: The T-DNA of this plasmid is schematically shown in Fig. 3 : Y_2 and the nucleotide sequence is depicted in SEQ ID N° 1.

² the mean of two samples independently measured (-0.2270 and -0.1963)

 $^{^{3}}$ The catalase activity in wild type SR1 tobacco leaves was set to 100 %.

Description of the transgene loci and production of hybrid plants

Locus X₁ harbours an inverted repeat about the right T-DNA border of construct pGVCHS287, carrying a neomycinphosphotransferase II (*nptII*) gene under the control of the Cauliflower mosaic virus 35S promoter (P35S) and the 3'signalling sequences of the Anthirrinum majus chalcone synthase gene (3'chs). The *nptII* genes are posttranscriptionally silenced and can trigger in trans silencing and methylation of homologous target genes (Van Houdt et al., 2000 a and b and Fig.2).

Locus Y_1 contains a single copy of the pGUSchsS T-DNA, containing a gus gene under the control of P35S and 3'chs (in transformant GUSchsS29) and shows normal levels of gus expression (Fig.2).

Locus Z₁ contains more than one copy of the pXD610 T-DNA, harbouring the gus gene under control of P35S and the 3'untranslated region (UTR) of the nopaline synthase gene (3'nos), (in plant LXD610-2) and shows normal gus expression (De Loose et al., 1995 and Fig.2).

Locus X₂ contains a single copy of both the pGUSchsS and pGUSchsAS T-DNA (in transformant GUSchsS+GUSchsAS 11) and triggers silencing in cis of the gus genes, but also in trans of (partially) homologous genes (Fig.4).

 X_1 and Z_1 hemizygous plants were obtained as hybrid progeny of the crossing of tobacco plants homozygous for locus X_1 (=Holo1; Van Houdt et al., 2000 a and b) and homozygous for locus Z_1 (=LXD610-2/9 De Loose et al., 1995) to wild type SR1 respectively. Y_1 hemizygous plants were obtained by crossing the hemizygous primary tobacco transformant GUSchsS29 to SR1 and selecting for the presence of locus Y_1 in the hybrid progeny. X_1Y_1 and Y_1Z_1 hemizygous plants are the hybrid progeny plants of the cross between Holo1 and GUSchsS29 and between GUSchsS29 and LXD610-2/9 respectively that are selected for the presence of Y_1 . X_1Z_1 hemizygous plants are the hybrid progeny of the cross between Holo1 and LXD610-2/9. $X_1Y_1Z_1$ hemizygous plants were obtained by crossing X_1Y_1 hemizygous plants to LXD610-2/9; as we only selected for the presence of Y_1 in the hybrid progeny both Y_1Z_1 and $X_1Y_1Z_1$ hemizygous plants were obtained.

Preparation of Agrobacteria and injection

10

20

25

30

The Agrobacteria C58C1Rif^R(pGV2260)(pGUSchsS)Cb^R,PPT^R or C58C1Rif^R(pMP90) (pPs35SCAT1S3chs)Gm^R,PPT^R were mainly grown as described by Kapila et al., 1997 except that the Agrobacteria were resuspended in MMA to a final OD₆₀₀ of 1.

Greenhouse grown plants of 10 to 15 cm in height were used. Half of the third top leaf was injected via the lower surface using a 5ml syringe while the leaf remained attached to the plant. The plants were kept in the greenhouse and 16 days after injection three to four discs of 11 mm in diameter were excised from the injected tissue for the preparation of a fresh protein extract to determine the catalase activity.

Enzymatic assays

5

10

Preparation of the protein extracts and GUS-activity measurements were done as previously described (Van Houdt et al., 2000 b). Preparation of the protein extracts for catalase-activity measurement and the spectrophotometric catalase-activity determination was done according to Champnongpol et al., 1996.

References

Van Houdt, H., Kovarik, A., Van Montagu, M., and Depicker, A. (2000 a). Cross-talk between posttranscriptionally silenced neomycin phosphotransferase II transgenes.

5 FEBS Lett. 467, 41-46.

Van Houdt, H., Kovarik, A., Van Montagu, M., and Depicker, A. (2000 b) Both sense and antisense RNAs are targets for the sense transgene-induced posttranscriptional silencing mechanism. *Mol. Gen. Genet.* 263, 995-1002.

De Loose, M., Danthinne, X., Van Bockstaele, E., Van Montagu, M. and Depicker, A.,

- 10 (1995) Different 5'leader sequences modulate β-glucuronidase accumulation levels in transgenic *Nicotiana tobacum* plants. *Euphytica* 85, 209-216.
 - Kapila, J., De Rycke, R., Van Montagu, M. and Angenon, G. (1997) An *Agrobacterium*-mediated transient gene expression system for intact leaves. *Plant Science* 122, 101-108.
- 15 Champnongpol, S., Willekens, H., Langebartels, C., Van Montagu, M., Inzé, D., and Van Camp, W. (1996) Transgenic tobacco with a reduced catalase activity develops necrotic lesions and induces pathogenesis-related expression under high light. *Plant J.* 10(3), 491-503.
- Thomas, C. L., Jones, L., Baulcombe, D.C. and Maule, A.J. (2001) Size constraints for targetting post-transcriptional gene silencing and for RNA-directed methylation in *Nicotiana benthamiana* using potato virus X vector. *Plant J.* 25(4), 417-425.
 - De Buck, S. and Depicker, A. (2001) Disruption of their palindromic arrangement leads to selective loss of DNA methylation in inversely repeated gus transgenes in Arabidopsis. *Mol. Gen. Genom.* 265, 1060-1068.

25

<u>Claims</u>

1. A method for obtaining efficient RNA silencing of a target gene comprising the introduction of a recombinant gene into a host that comprises a silenced locus and a target gene whereby said recombinant gene comprises a region that is homologous with said silenced locus and whereby said target gene has homology with said recombinant gene but has no significant homology with said silenced locus.

2. A method according to claim 1 wherein said host is a plant or plant cell.

5

- 3. A method according to claims 1 or 2 to obtain high throughput gene silencing.
- 4. A plant or plant cell comprising a silenced target gene obtainable by a method according to claims 1 or 2.
 - 5. A method according to claims 1 or 2 wherein said RNA silencing of the target gene is obtained in more than 95% of the hosts.
 - 6. A method according to claims 1 or 2 wherein RNA silencing of the target gene is obtained in more than 85% of the hosts.
- 7. A method according to claims 1 or 2 wherein said RNA silencing of the target gene occurs at an efficiency of more than 95 % as compared to the level of the unsilenced expression of the target gene.
- 8. A method according to claims 1 or 2 wherein said RNA silencing of the target gene occurs at an efficiency of more than 85 % as compared to the level of the unsilenced expression of the target gene.

WO 03/031632

PCT/EP02/11188

1/5

Figure 1

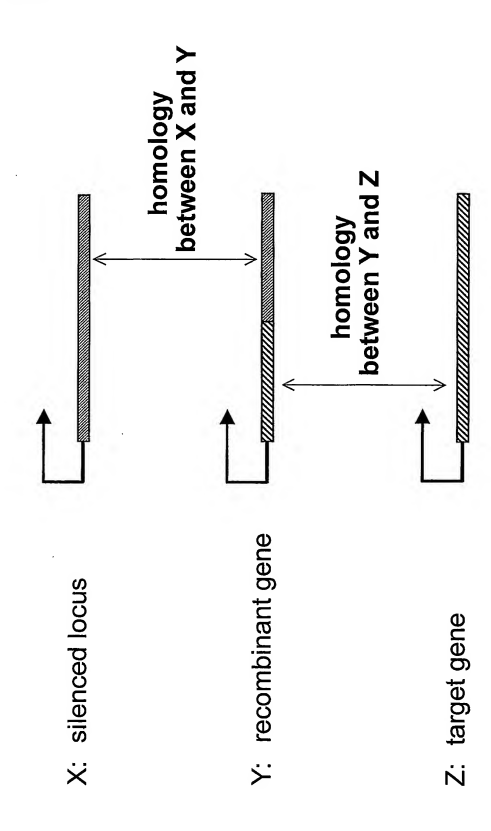


Figure 2

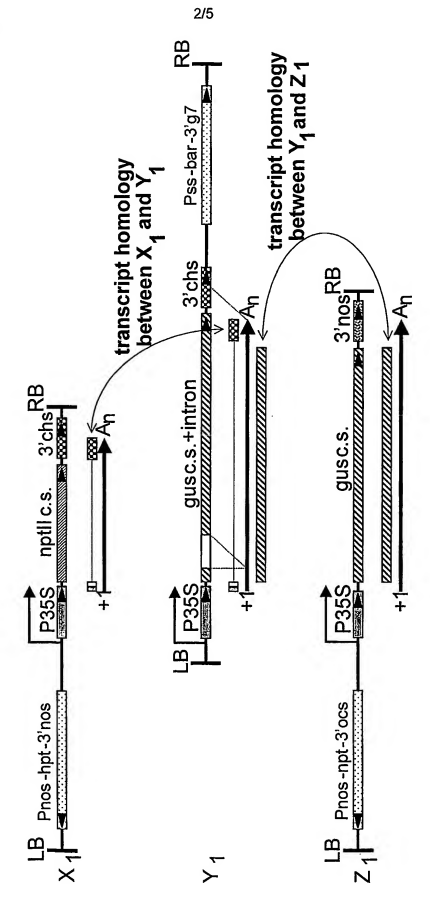
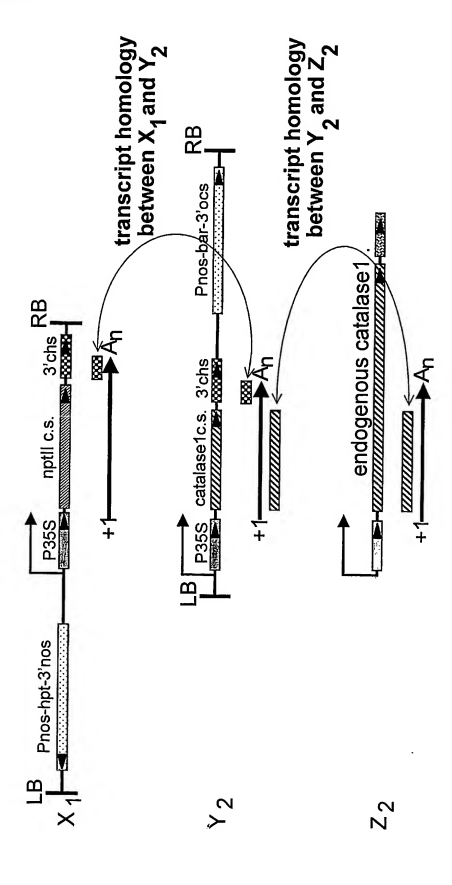
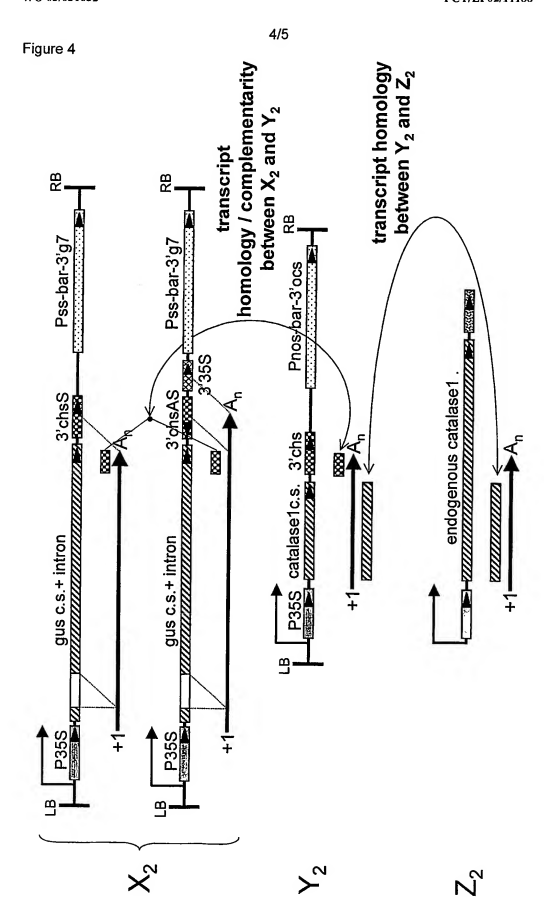


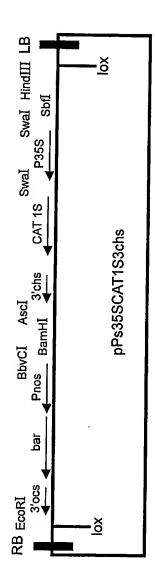
Figure 3





5/5

Figure 5



SEQUENCE LISTING

<110> VLAAMS INTERUNIVERSITAIR INSTITUUT VOOR BIOTECHNOLOGIE VZW <120> An efficient system for RNA silencing <130> ADP/Dom/V097 <150> EP01203760.2 <151> 2001-10-05 <160> 1 <170> PatentIn version 3.1 <210> 1 <211> 10635 <212> DNA <213> Artificial Sequence <220> <223> pPs35SCAT1S3chs agattcgaag ctcggtcccg tgggtgttct gtcgtctcgt tgtacaacga aatccattcc 60 cattccgcgc tcaagatggc ttcccctcgg cagttcatca gggctaaatc aatctagccg 120 acttgtccgg tgaaatgggc tgcactccaa cagaaacaat caaacaaaca tacacagcga 180 cttattcaca cgcgacaaat tacaacggta tatatcctgc cagtactcgg ccgtcgaata 240 acttcgtata atgtatgcta tacgaagtta tgaattcgcg ctctatcata gatgtcgcta 300 taaacctatt cagcacaata tattgttttc attttaatat tgtacatata agtagtaggg 360 tacaatcagt aaattgaacg gagaatatta ttcataaaaa tacgatagta acgggtgata 420 tattcattag aatgaaccga aaccggcggt aaggatctga gctacacatg ctcaggtttt 480 ttacaacgtg cacaacagaa ttgaaagcaa atatcatgcg atcataggcg tctcgcatat 540 ctcattaaag Cagctggaag atttgatgga tcctcatcag atctcggtga cgggcaggac 600 cggacggggc ggtaccggca ggctgaagtc cagctgccag aaacccacgt catgccagtt 660 cccgtgcttg aagccggccg cccgcagcat gccgcggggg gcatatccga gcgcctcgtg 720 catgcgcacg ctcgggtcgt tgggcagccc gatgacagcg accacgctct tgaagccctg 780 tgcctccagg gacttcagca ggtgggtgta gagcgtggag cccagtcccg tccgctggtg 840 gcggggggag acgtacacgg tcgactcggc cgtccagtcg taggcgttgc gtgccttcca 900

ggggcccgcg taggcgatgc cggcgacctc gccgtccacc tcggcgacga gccagggata 960 gcgctcccgc agacggacga ggtcgtccgt ccactcctgc ggttcctgcg gctcggtacg 1020 gaagttgacc gtgcttgtct cgatgtagtg gttgacgatg gtgcagaccg ccggcatgtc 1080 cgcctcggtg gcacggcgga tgtcggccgg gcgtcgttct gggctcatgg tagatctgtt 1140 taaacgttaa cggattgaga gtgaatatga gactctaatt ggataccgag gggaatttat 1200 ggaacgtcag tggagcattt ttgacaagaa atatttgcta gctgatagtg accttaggcg 1260 acttttgaac gcgcaataat ggtttctgac gtatgtgctt agctcattaa actccagaaa 1320 cccgcggctg agtggctcct tcaatcgttg cggttctgtc agttccaaac gtaaaacggc 1380 ttgtcccgcg tcatcggcgg gggtcataac gtgactccct taattctccg ctcatgatca 1440 agctacctca gcaggatccg gcgcgccatg gtcgataaga aaaggcaatt tgtagatgtt 1500 aattcataac atctcctcca tgacttaaaa aacttgcaaa agatttatat agaaatactt 1560 aaatattttg actaaaaaaa aaaaaaaaaa aacacacaca taaaccaaca aataacataa 1620 attattttta tatagccttt atttcaatga tcacaacgaa acaatacaag tacaaagcgt 1680 tacaagagag aaatcgccaa tatagctcac atgcagcaca catcacaata ataggtaacc 1740 atgtccactt ttttattacg gaaataagaa aataacccaa cccccgtacc cgggttcata 1800 tgcttggtct cacattaagc ctagaagcta gcttttgacc cagagatttg tcagcctgag 1860 accagtatga gatccaaatg ctgcggatct cataagtgat acgaggatca gacaaggtct 1920 ccacccaccg acgaataaag cgttcttgcc tgtctggtgt gaatgagcgg tacctttctc 1980 ctggttgctt gaaattgttc tctttctgaa tgacacactt ctcgcgtttg ccagtgcaca 2040 ttgtagaagg aataggatac ttctcagcat ggcgaacagg atcatacctt gaagggaagt 2100 agtcgatctc ctcatccctg tgcataaaat tcatggagcc atcgtagtga ttgttgtgat 2160 gagcgcattt tggagcatta gcaggtagtt gcaaatagtt tggtccaagt cgatacctct 2220 gggtatcaga gtaggagaaa atacgagttt gaagcatctt atcatctgag taataaaccc 2280 ctggaacaac aatagaaggg cagaaagcta gctgctcatt ctcattagag aagttatcaa 2340 tgttcttgtt cagaactaat cttcccaccg gctgcaaagg caagatatcc tctggccaag 2400 tttttgtcac atcaagtgga tcaaaatcaa atctgtcttc atgatctgga tccatagtcc 2460 ccgggcagtg ggcgatttga tttaaatctc tagaatagta aattgtaatg ttgtttgttg 2520 tttgttttgt tgtggtaatt gttgtaaaaa tacggatcgt cctgcagtcc tctccaaatg 2580 aaatgaactt ccttatatag aggaagggtc ttgcgaagga tagtggggatt gtgcgtcatc 2640 ccttacgtca gtggagatat cacatcaatc cacttgcttt gaagacgtgg ttggaacgtc 2700 ttctttttcc acgatgctcc tcgtgggtgg gggtccatct ttgggaccac tgtcggcaga 2760 ggcatcttga acgatagcct ttcctttatc gcaatgatgg catttgtagg tgccaccttc 2820 cttttctact gtccttttga tgaagtgaca gatagctggg caatggaatc cgaggaggtt 2880 tcccgatatt accctttgtt gaaaagtctc aatagccctt tggtcttctg agactgtatc 2940 tttgatattc ttggagtaga cgagagtgtc gtgctccacc atgttgacga agattttctt 3000 cttgtcattg agtcgtaaaa gactctgtat gaactgttcg ccagtcttca cggcgagttc 3060 tgttagatcc tcgatctgaa tttttgactc catggccttt gattcagtag gaactacttt 3120 cttagagact ccaatctcta ttacttgcct tggtttatga agcaagcctt gaatcgtcca 3180

tactggaata	gtacttctga	tcttgagaaa	tatatctttc	tctgtgttct	tgatgcagtt	3240
agtcctgaat	cttttgactg	catctttaac	cttcttggga	aggtatttga	tctcctggag	3300
attattacto	gggtagatcg	tcttgatgag	acctgccgcg	taggcctctc	taaccatctg	3360
tgggtcagca	ttctttctga	aattgaagag	gctaatcttc	tcattatcgg	tggtgaacat	3420
ggtatcgtca	ccttctccgt	cgaactttct	tcctagatcg	tagagataga	gaaagtcgtc	3480
catggtgatc	tccggggcaa	aggagatctc	tagagtcgag	atttaaatcc	taaatcctgc	3540
aggaagctta	ccggtataac	ttcgtatagc	atacattata	cgaagttatc	catggagcca	3600
tttacaattg	aatatatcct	gccgccgctg	ccgctttgca	cccggtggag	cttgcatgtt	3660
ggtttctacg	cagaactgag	ccggttaggc	agataatttc	cattgagaac	tgagccatgt	3720
gcaccttccc	cccaacacgg	tgagcgacgg	ggcaacggag	tgatccacat	gggactttta	3780
aacatcatcc	gtcggatggc	gttgcgagag	aagcagtcga	tccgtgagat	cagccgacgc	3840
accgggcagg	cgcgcaacac	gatcgcaaag	tatttgaacg	caggtacaat	cgagccgacg	3900
ttcacggtac	cggaacgacc	aagcaagcta	gcttagtaaa	gccctcgcta	gattttaatg	3960
cggatgttgc	gattacttcg	ccaactattg	cgataacaag	aaaaagccag	cctttcatga	4020
tatatctccc	aatttgtgta	gggcttatta	tgcacgctta	aaaataataa	aagcagactt	4080
gacctgatag	tttggctgtg	agcaattatg	tgcttagtgc	atctaacgct	tgagttaagc	4140
cgcgccgcga	agcggcgtcg	gcttgaacga	attgttagac	attatttgcc	gactaccttg	4200
gtgatctcgc	ctttcacgta	gtggacaaat	tcttccaact	gatctgcgcg	cgaggccaag	4260
cgatcttctt	cttgtccaag	ataagcctgt	ctagcttcaa	gtatgacggg	ctgatactgg	4320
gccggcaggc	gctccattgc	ccagtcggca	gcgacatcct	tcggcgcgat	tttgccggtt	4380
actgcgctgt	accaaatgcg	ggacaacgta	agcactacat	ttcgctcatc	gccagcccag	4440
tcgggcggcg	agttccatag	cgttaaggtt	tcatttagcg	cctcaaatag	atcctgttca	4500
ggaaccggat	caaagagttc	ctccgccgct	ggacctacca	aggcaacgct	atgttctctt	4560
gcttttgtca	gcaagatagc	cagatcaatg	tcgatcgtgg	ctggctcgaa	gatacctgca	4620
agaatgtcat	tgcgctgcca	ttctccaaat	tgcagttcgc	gcttagctgg	ataacgccac	4680
ggaatgatgt	cgtcgtgcac	aacaatggtg	acttctacag	cgcggagaat	ctcgctctct	4740
ccaggggaag	ccgaagtttc	caaaaggtcg	ttgatcaaag	ctcgccgcgt	tgtttcatca	4800
agccttacgg	tcaccgtaac	cagcaaatca	atatcactgt	gtggcttcag	gccgccatcc	4860
actgcggagc	cgtacaaatg	tacggccagc	aacgtcggtt	cgagatggcg	ctcgatgacg	4920
ccaactacct	ctgatagttg	agtcgatact	tcggcgatca	ccgcttccct	catgatgttt	4980
aactttgttt	tagggcgact	gccctgctgc	gtaacatcgt	tgctgctcca	taacatcaaa	5040
catcgaccca	cggcgtaacg	cgcttgctgc	ttggatgccc	gaggcataga	ctgtacccca	5100
		catgaaaacc				5160
		gcgtgagcgc				5220
		ggttcgtgcc				5280
		agtcgaggca				5340
gtttcggtct	ccacgcatcg	tcaggcattg	gcggccttgc	tgttcttcta	cggcaagtgc	5400

tgtgcacgga	tctgccctgg	cttcaggaga	tcggaagacc	tcggccgtcc	gggcgcttgc	5460
cggtggtgct	gaccccggat	gaagtggttc	gcatcctcgg	ttttctggaa	ggcgagcatc	5520
gtttgttcgc	ccagcttctg	tatggaacgg	gcatgcggat	cagtgagggt	ttgcaactgc	5580
gggtcaagga	tctggatttc	gatcacggca	cgatcatcgt	gcgggagggc	aagggctcca	5640
aggatcgggc	cttgatgtta	cccgagagct	tggcacccag	cctgcgcgag	cagggatcga	5700
tccaacccct	ccgctgctat	agtgcagtcg	gcttctgacg	ttcagtgcag	ccgtcttctg	5760
aaaacgacat	gtcgcacaag	tcctaagtta	cgcgacaggc	tgccgccctg	cccttttcct	5820
ggcgttttct	tgtcgcgtgt	tttagtcgca	taaagtagaa	tacttgcgac	tagaaccgga	5880
gacattacgc	catgaacaag	agcgccgccg	ctggcctgct	gggctatgcc	cgcgtcagca	5940
ccgacgacca	ggacttgacc	aaccaacggg	ccgaactgca	cgcggccggc	tgcaccaagc	6000
tgttttccga	gaagatcacc	ggcaccaggc	gcgaccgccc	ggagctggcc	aggatgcttg	6060
accacctacg	ccctggcgac	gttgtgacag	tgaccaggct	agaccgcctg	gcccgcagca	6120
cccgcgacct	actggacatt	gccgagcgca	tccaggaggc	cggcgcgggc	ctgcgtagcc	6180
tggcagagcc	gtgggccgac	accaccacgc	cggccggccg	catggtgttg	accgtgttcg	6240
ccggcattgc	cgagttcgag	cgttccctaa	tcatcgaccg	cacccggagc	gggcgcgagg	6300
ccgccaaggc	ccgaggcgtg	aagtttggcc	cccgccctac	cctcaccccg	gcacagatcg	6360
cgcacgcccg	cgagctgatc	gaccaggaag	gccgcaccgt	gaaagaggcg	gctgcactgc	6420
ttggcgtgca	tcgctcgacc	ctgtaccgcg	cacttgagcg	cagcgaggaa	gtgacgccca	6480
ccgaggccag	gcggcgcggt	gccttccgtg	aggacgcatt	gaccgaggcc	gacgccctgg	6540
cggccgccga	gaatgaacgc	caagaggaac	aagcatgaaa	ccgcaccagg	acggccagga	6600
cgaaccgttt	ttcattaccg	aagagatcga	ggcggagatg	atcgcggccg	ggtacgtgtt	6660
cgagccgccc	gcgcacgtct	caaccgtgcg	gctgcatgaa	atcctggccg	gtttgtctga	6720
tgccaagctg	gcggcctggc	cggccagctt	ggccgctgaa	gaaaccgagc	gccgccgtct	6780
aaaaaggtga	tgtgtatttg	agtaaaacag	cttgcgtcat	gcggtcgctg	cgtatatgat	6840
gcgatgagta	aataaacaaa	tacgcaaggg	gaacgcatga	aggttatcgc	tgtacttaac	6900
cagaaaggcg	ggtcaggcaa	gacgaccatc	gcaacccatc	tagcccgcgc	cctgcaactc	6960
gccggggccg	atgttctgtt	agtcgattcc	gatccccagg	gcagtgcccg	cgattgggcg	7020
				accgcccgac		7080
gacgtgaagg	ccatcggccg	gcgcgacttc	gtagtgatcg	acggagcgcc	ccaggcggcg	7140
gacttggctg	tgtccgcgat	caaggcagcc	gacttcgtgc	tgattccggt	gcagccaagc	7200
ccttacgaca	tatgggccac	cgccgacctg	gtggagctgg	ttaagcagcg	cattgaggtc	7260
				cgatcaaagg		7320
				ccattcttga		7380
				caaccgttct		7440
cccgagggcg	acgctgcccg	cgaggtccag	gcgctggccg	ctgaaattaa	atcaaaactc	7500
				aaacacgcta		7560
				cctggcagac	_	7620
tgaagcgggt	caactttcag	ttgccggcgg	aggatcacac	caagctgaag	atgtacgcgg	7680

tacgccaagg	caagaccatt	accgagctgc	tatctgaata	catcgcgcag	ctaccagagt	7740
aaatgagcaa	atgaataaat	gagtagatga	attttagcgg	ctaaaggagg	cggcatggaa	7800
aatcaagaac	aaccaggcac	cgacgccgtg	gaatgcccca	tgtgtggagg	aacgggcggt	7860
tggccaggcg	taagcggctg	ggttgtctgc	cggccctgca	atggcactgg	aacccccaag	7920
cccgaggaat	cggcgtgacg	gtcgcaaacc	atccggcccg	gtacaaatcg	gcgcggcgct	7980
gggtgatgac	ctggtggaga	agttgaaggc	cgcgcaggcc	gcccagcggc	aacgcatcga	8040
ggcagaagca	cgccccggtg	aatcgtggca	agcggccgct	gatcgaatcc	gcaaagaatc	8100
ccggcaaccg	ccggcagccg	gtgcgccgtc	gattaggaag	ccgcccaagg	gcgacgagca	8160
accagatttt	ttcgttccga	tgctctatga	cgtgggcacc	cgcgatagtc	gcagcatcat	8220
ggacgtggcc	gttttccgtc	tgtcgaagcg	tgaccgacga	gctggcgagg	tgatccgcta	8280
cgagcttcca	gacgggcacg	tagaggtttc	cgcagggccg	gccggcatgg	ccagtgtgtg	8340
ggattacgac	ctggtactga	tggcggtttc	ccatctaacc	gaatccatga	accgataccg	8400
ggaagggaag	ggagacaagc	ccggccgcgt	gttccgtcca	cacgttgcgg	acgtactcaa	8460
gttctgccgg	cgagccgatg	gcggaaagca	gaaagacgac	ctggtagaaa	cctgcattcg	8520
gttaaacacc	acgcacgttg	ccatgcagcg	tacgaagaag	gccaagaacg	gccgcctggt	8580
gacggtatcc	gagggtgaag	ccttgattag	ccgctacaag	atcgtaaaga	gcgaaaccgg	8640
gcggccggag	tacatcgaga	tcgagctagc	tgattggatg	taccgcgaga	tcacagaagg	8700
caagaacccg	gacgtgctga	cggttcaccc	cgattacttt	ttgatcgatc	ccggcatcgg	8760
ccgttttctc	taccgcctgg	cacgccgcgc	cgcaggcaag	gcagaagcca	gatggttgtt	8820
caagacgatc	tacgaacgca	gtggcagcgc	cggagagttc	aagaagttct	gtttcaccgt	8880
gcgcaagctg	atcgggtcaa	atgacctgcc	ggagtacgat	ttgaaggagg	aggcggggca	8940
ggctggcccg	atcctagtca	tgcgctaccg	caacctgatc	gagggcgaag	catccgccgg	9000
ttcctaatgt	acggagcaga	tgctagggca	aattgcccta	gcaggggaaa	aaggtcgaaa	9060
aggtctcttt	cctgtggata	gcacgtacat	tgggaaccca	aagccgtaca	ttgggaaccg	9120
gaacccgtac	attgggaacc	caaagccgta	cattgggaac	cggtcacaca	tgtaagtgac	9180
tgatataaaa	gagaaaaaag	gcgattttc	cgcctaaaac	tctttaaaac	ttattaaaac	9240
tcttaaaacc	cgcctggcct	gtgcataact	gtctggccag	cgcacagccg	aagagctgca	9300
	acccttcggt					9360
cgcggccgct		-				9420
acaagccgcg						9480
gtttcggtga						9540
gtctgtaagc						9600
ggtgtcgggg						9660
ctatgcggca						9720
cagatgcgta						9780
gctgcgctcg						9840
gttatccaca	gaatcagggg	ataacgcagg	aaagaacatg	tgagcaaaag	gccagcaaaa	9900

ggccaggaac cgtaaa	aagg ccgcgttgct	ggcgtttttc	cataggctcc	gccccctga	9960
cgagcatcac aaaaat	cgac gctcaagtca	gaggtggcga	aacccgacag	gactataaag	10020
ataccaggcg tttcccc	cctg gaagctccct	cgtgcgctct	cctgttccga	ccctgccgct	10080
taccggatac ctgtccg	gcct ttctcccttc	gggaagcgtg	gcgctttctc	atagctcacg	10140
ctgtaggtat ctcagt	tcgg tgtaggtcgt	tcgctccaag	ctgggctgtg	tgcacgaacc	10200
ccccgttcag cccgaco	gct gcgccttatc	cggtaactat	cgtcttgagt	ccaacccggt	10260
aagacacgac ttatcgo	ccac tggcagcagc	cactggtaac	aggattagca	gagcgaggta	10320
tgtaggcggt gctacag	gagt tcttgaagtg	gtggcctaac	tacggctaca	ctagaaggac	10380
agtatttggt atctgcg	yctc tgctgaagcc	agttaccttc	ggaaaaagag	ttggtagctc	10440
ttgatccggc aaacaaa	acca ccgctggtag	cggtggtttt	tttgtttgca	agcagcagat	10500
tacgcgcaga aaaaaag	ggat ctcaagaaga	tccggaaaac	gcaagcgcaa	agagaaagca	10560
ggtagcttgc agtgggd	tta catggcgata	gctagactgg	gcggttttat	ggacagcaag	10620
cgaaccggaa ttgcc					10635

INTERNATIONAL SEARCH REPORT

Intervious Application No PCT/EP 02/11188

			C1/EP UZ/11188
A. CLASSI IPC 7	IFICATION OF SUBJECT MATTER C12N15/82		
According to	o International Patent Classification (IPC) or to both national classifica	ation and IPC	
B. FIELDS	SEARCHED		
Minimum do	ocumentation searched (classification system followed by classification ${\tt C12N}$	on symbols)	
	tion searched other than minimum documentation to the extent that so	į	
	lata base consulted during the international search (name of data bas ta, EPO-Internal, PAJ, BIOSIS, EMBAS	•	arch (erms used)
C DOCUME	ENTS CONSIDERED TO BE RELEVANT		
Category °	Citation of document, with indication, where appropriate, of the rela	munet nonconno	Delevent to eleim No.
Calegory	Citation of occurrent, was muscation, whose appropriate, or me resc	evant passages	Relevant to claim No.
A	VAN HOUDT H ET AL: "Cross-talk b posttranscriptionally silenced ne phosphotransferase II transgenes"	eomycin	1-8
	FEBS LETTERS, ELSEVIER SCIENCE PU AMSTERDAM, NL, vol. 467, no. 1,		
	4 February 2000 (2000-02-04), pag XP004260919 the whole document	jes 41-46,	
A	WO 99 53050 A (WANG MING BO ;COMM IND RES ORG (AU); GRAHAM MICHAEL		1-8
	21 October 1999 (1999-10-21) cited in the application the whole document	WATINE)	·
		-/	
X Furth	her documents are listed in the continuation of box C.	X Patent family me	mbers are listed in annex.
Special car	ategories of cited documents:	*T* later document publish	ed after the International filing date
consid	ent defining the general state of the art which is not dered to be of particular relevance document but published on or after the international	cited to understand the invention	t in conflict with the application but e principle or theory underlying the
filing d "L" docume which i	ant which may throw doubts on priority ctaim(s) or	cannot be considered involve an inventive s	relevance; the claimed invention novel or cannot be considered to tep when the document is taken alone
citation	n or other special reason (as specified) ent referring to an oral disclosure, use, exhibition or	cannot be considered document is combine	relevance; the claimed invention to involve an inventive step when the d with one or more other such docu- ion being obvious to a person skilled
"P" docume	ent published prior to the international filing date but	in the art. *&" document member of t	
Date of the	actual completion of the International search	Date of mailing of the	International search report
1:	3 December 2002	20/12/200	2
Name and n	malling address of the ISA European Patent Office, P.B. 5818 Patentlaan 2	Authorized officer	
	NL – 2280 HV Rijswijk Tel. (+31–70) 340–2040, Tx. 31 651 epo nl, Fax: (+31–70) 340–3016	Kalsner,	I

INTERNATIONAL SEARCH REPORT

Interaction No PCT/EP 02/11188

Category* Citation of document, with indication,where appropriate, of the rethrent passages A MO 99 61632 A (NAP JAN PETER HENDRIK ;CPRO DLO (NL); STIEKEMA WILLEM JOHANNES (NL) 2 December 1999 (1999–12–02) cited in the application the whole document			PCT/EP 02/11188		
A WO 99 61632 A (NAP JAN PETER HENDRIK ;CPRO 1-8 DLO (NL); STIEKEMA WILLEM JOHANNES (NL) 2 December 1999 (1999-12-02) cited in the application					
Cited in the application	Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.		
	Category °	Citation of document, with indication, where appropriate, of the relevant passages WO 99 61632 A (NAP JAN PETER HENDRIK; CPRO DLO (NL); STIEKEMA WILLEM JOHANNES (NL) 2 December 1999 (1999–12–02) cited in the application			

INTERNATIONAL SEARCH REPORT

Information on patent family members

Intertional Application No PCT/EP 02/11188

Patent document cited in search report		Publication date		Patent family member(s)	Publication date
WO 9953050	A	21-10-1999	AU CA CN EP WO JP	2951499 A 2325344 A1 1306571 T 1068311 A1 9953050 A1 2002511258 T	01-11-1999 21-10-1999 01-08-2001 17-01-2001 21-10-1999 16-04-2002
WO 9961632	A	02-12-1999	EP AU WO	0959133 A1 4172599, A 9961632 A1	24-11-1999 13-12-1999 02-12-1999